Atty. Dkt. No. GXI-10

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**PATENT** 

## **IN THE CLAIMS**

In the claims, please cancel Claims 1 through 25. Please add Claims 26 through 47. A clean version of all of the claims is included below. Please add the claims:

26. (NEW) A method for eliminating redundant sequences that are common between two samples, the method comprising of the steps:

isolating RNA strands from a first sample;
isolating RNA strands from a second sample;
generating cDNA strands from the RNA strands from the first sample;
mixing the cDNA strands of the first sample with the RNA strands from the
second sample, the cDNA strands and RNA strands with common sequences
hybridizing to form cDNA/RNA compliments; and
degrading the compliments.

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- 27. (NEW) The method of Claim 26, wherein the step of generating cDNA strands for the RNA strands from the first sample comprises performing RT-PCR.
- 28. (NEW) The method of Claim 27, wherein the step of performing RT-PCR utilizes radioactive labeled to anchored oligo dT primers.
- 29. (NEW) The method of Claim 26, wherein the first sample is a healthy tissue and the second sample is a diseased tissue.
- 30. (NEW) The method of Claim 26, wherein the first sample is a diseased tissue and the second sample is a healthy tissue.
- 31. (NEW) The method of Claim 26, further comprising: amplifying the unhybridized cDNA strands using PCR.
- 32. (NEW) The method of Claim 26, further comprising: producing a second set of cDNA strands from unhybridized RNA strands.
- 33. (NEW) The method of Claim 32, further comprising: amplifying the second set of cDNA strands using PCR.

- 34. (NEW) The method of Claim 26, wherein the step of degrading compliments is performed with an Exonuclease III enzyme.
- 35. (NEW) The method of Claim 26, wherein the step of degrading compliments is performed with an Exonuclease IV enzyme.
- 36. (NEW) The method of Claim 26, further comprising: displaying the unhybridized cDNAs or RNAs
- 37. (NEW) The method of Claim 36, wherein the step of displaying comprises using electrophoresis.
- 38. (NEW) The method of Claim 26, further comprising:
  reading the unhybridized cDNA strands or RNA strands with a photographic plate.
- 39. (NEW) The method of Claim 26, wherein the first and second samples are selected from a group comprising cells, tissues, pathogens, plants, and animals.
- 40. (NEW) The method of Claim 26, wherein the first and second sample are differentiated due to a diseased state, developmental, change, or induced by an external or internal stimulus.
- 41. (NEW) A method for determining differences between a first sample of cDNA strands and a second sample of RNA strands, the method comprising of the steps:

  mixing the first sample of cDNA strands with the second sample of RNA strands, the cDNA strands and RNA strands with common sequences hybridizing to form cDNA/RNA compliments; and degrading the cDNA/RNA compliments.

- 42. (NEW) The method of Claim 41, wherein the first sample of cDNA strands are produced from a third sample of RNA strands using RT-PCR utilizes radioactive labeled to anchored oligo dT primers.
- 43. (NEW) The method of Claim 41, further comprising: amplifying unhybridized cDNA strands using PCR.
- 44. (NEW) The method of Claim 41, further comprising: producing a further set of cDNA strands from unhybridized RNA strands.
- 45. (NEW) The method of Claim 44, further comprising: amplifying the further set of cDNA strands using PCR.
- 46. (NEW) The method of Claim 41, wherein the step of degrading compliments is performed with an Exonuclease III enzyme.
- 47. (NEW) The method of Claim 41, wherein the step of degrading compliments is performed with an Exonuclease IV enzyme.